

Dental data challenge the ubiquitous presence of *Homo* in the Cradle of Humankind

Clément Zanolli^{a,1}, Thomas W. Davies^{b,c}, Renaud Joannes-Boyau^{d,e}, Amélie Beaudet^{f,g,h}, Laurent Bruxelles^{g,ij}, Frikkie de Beer^{k,i}, Jakobus Hoffman^k, Jean-Jacques Hublin^b, Kudakwashe Jakata^m, Lazarus Kgasi^{e,n}, Ottmar Kullmer^{o,p}, Roberto Macchiarelli^{q,r}, Lei Pan^{s,t,2}, Friedemann Schrenk^{o,p}, Frédéric Santos^a, Dominic Stratford^g, Mirriam Tawaneⁿ, Francis Thackeray^m, Song Xing^{s,t}, Bernhard Zipfel^m, and Matthew M. Skinner^{b,c,u}

Edited by Lucas Delezene, University of Arkansas Fayetteville, Fayetteville, AR; received June 17, 2021; accepted April 13, 2022 by Editorial Board Member Richard G. Klein

The origins of Homo, as well as the diversity and biogeographic distribution of early Homo species, remain critical outstanding issues in paleoanthropology. Debates about the recognition of early Homo, first appearance dates, and taxonomic diversity within Homo are particularly important for determining the role that southern African taxa may have played in the origins of the genus. The correct identification of Homo remains also has implications for reconstructing phylogenetic relationships between species of Australopithecus and Paranthropus, and the links between early Homo species and Homo erectus. We use microcomputed tomography and landmark-free deformationbased three-dimensional geometric morphometrics to extract taxonomically informative data from the internal structure of postcanine teeth attributed to Early Pleistocene Homo in the southern African hominin-bearing sites of Sterkfontein, Swartkrans, Drimolen, and Kromdraai B. Our results indicate that, from our sample of 23 specimens, only 4 are unambiguously attributed to Homo, 3 of them coming from Swartkrans member 1 (SK 27, SK 847, and SKX 21204) and 1 from Sterkfontein (Sts 9). Three other specimens from Sterkfontein (StW 80 and 81, SE 1508, and StW 669) approximate the Homo condition in terms of overall enamel-dentine junction shape, but retain Australopithecus-like dental traits, and their generic status remains unclear. The other specimens, including SK 15, present a dominant australopith dental signature. In light of these results, previous dietary and ecological interpretations can be reevaluated, showing that the geochemical signal of one tooth from Kromdraai (KB 5223) and two from Swartkrans (SK 96 and SKX 268) is consistent with that of australopiths.

early Homo | taxonomic assessment | dental structure | geometric morphometrics

The South African cave sites of the Cradle of Humankind (Gauteng Province), declared a World Heritage area by the United Nations Educational, Scientific, and Cultural Organization, have yielded hundreds of Pliocene-Pleistocene (Gelasian-Calabrian, https:// stratigraphy.org) hominin remains assigned to the genera Australopithecus, Paranthropus, and Homo. At Sterkfontein, specimens variously assigned to Australopithecus, Paranthropus robustus, and early Homo have been found in western breccias equivalent in age to post-member 4 infills (member 5 and the StW 53 infill) (1, 2). Currently available dates for the StW 53 infill and member 5 deposits suggest that the genera associated with these units existed in close temporal proximity on the southern African landscape during the Late Pliocene to Early Pleistocene (3), even if a better chronological framework is needed to accurately compare specimens from the various sites. Additionally, the identification of specimens belonging to early Homo that are penecontemporaneous with Australopithecus and/or Paranthropus at sites such as Kromdraai B, Drimolen, and Swartkrans has led to interpretations of a widespread presence of early Homo throughout the Cradle of Humankind since ~2.5 Ma (see debates about the geochronology of southern African hominin sites in refs. 4-7). However, the attribution of a number of these specimens to Homo has been questioned. In this study, we analyze the dentine shape of postcanine tooth crowns, which has demonstrated a strong taxonomic signal at the generic level. We reexamine the taxonomic attribution of the majority of specimens previously considered to represent early Homo, and test the hypothesis of a ubiquitous presence of early Homo in the Early Pleistocene sites of southern Africa.

The first species described for *Australopithecus*, *Australopithecus africanus*, was erected based on the Late Pliocene juvenile skull Taung 1 from the eponymous site (8), and the hypodigm was later supplemented by specimens from Sterkfontein, Makapansgat, and Gladysvale (9). *Australopithecus prometheus* was first described on the basis of the

Significance

Identifying the earliest members of the genus *Homo* is crucial for understanding when and where selective pressures resulted in its emergence from a Plio-Pleistocene hominin taxon. Our revision of a large part of the dental fossil record from southern Africa provides evidence suggesting a paucity of Homo remains and indicates increased levels of dental variation in australopith taxa. Results of the Ba/Ca, Sr/Ca, and elemental mapping of enamel and dentine also indicate that some of the purported Homo specimens show a paleoecological signal similar to that of the australopiths.

Author contributions: C.Z. and M.M.S. designed research; C.Z. and M.M.S. performed research; C.Z., T.W.D., R.J.-B., A.B., and M.M.S. analyzed data; C.Z., T.W.D., R.J.-B., A.B., L.B., F.d.B., J.H., J.-J.H., K.J., L.K., O.K., R.M., L.P., F. Schrenk, F. Santos, D.S., M.T., F.T., S.X., B.Z., and M.M.S. wrote the paper; L.B. contributed to the revision of stratigraphy and chronology of the sites; F.d.B. contributed to the 3D scanning and elaboration of the microCT scans; J.H. and K.J. contributed to the microCT scanning of the fossil specimens and elaboration of the 3D data; J.H., L.K., O.K., R.M., L.P., F. Schrenk, D.S., M.T., F.T., S.X., and B.Z. discussed the implications of the findings; and F. Santos contributed to the statistical analyses.

The authors declare no competing interest.

This article is a PNAS Direct Submission. L.D. is a guest editor invited by the Editorial Board.

Copyright © 2022 the Author(s). Published by PNAS. This article is distributed under Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND).

¹To whom correspondence may be addressed. Email: clement.zanolli@gmail.com.

²Deceased May 2020.

This article contains supporting information online at http://www.pnas.org/lookup/suppl/doi:10.1073/pnas. 2111212119/-/DCSupplemental.

Published July 5, 2022.

parietooccipital portion of braincase MLD 1 from Makapansgat (10). It was also recognized in the Late Pliocene deposits of Sterkfontein (11). The Early Pleistocene species *Australopithecus sediba* was identified at Malapa and described as showing *Homo*-like features even if the postcanine teeth exhibit an *Australopithecus* morphology, including birooted lower third premolars and the presence of a marked protostylid on the lower molars (12). The type species of *Paranthropus, P. robustus*, was described on the basis of the Early Pleistocene partial skull TM 1517 from Kromdraai (13) and was recognized at several other sites such as Drimolen, Cooper's Cave, Gondolin, Sterkfontein, and Swartkrans (9).

While less plentiful, remains attributed to early Homo have been described in four sites of the Cradle of Humankind: Swartkrans, Kromdraai B, Sterkfontein, and Drimolen. The first fossil from southern Africa attributed to early Homo was the Early Pleistocene mandible SK 15 from Swartkrans member 2, dated to ~1.4 Ma (14). It was originally attributed to Telanthropus capensis (15) and was later assigned to Homo erectus/ Homo ergaster (16), but it was recently suggested to belong to Australopithecus (17). Additional specimens, including SK 27, SK 847, and SKX 21204 from the 2.2 Ma to 1.8 Ma Swartkrans member 1 (7), and the isolated teeth SKX 257 and 258 and SKX 267 through 269 from the chronologically younger member 2, are generally recognized as early Homo (18-20). Several specimens from this site, including SK 47, SK 843, SK 846, and SKX 4446, were previously attributed to Homo (21, 22), but further analyses suggested that they belong to the P. robustus hypodigm (23, 24).

The associated dentition of KB 5223 from Kromdraai B, chronologically assessed to the beginning of the Early Pleistocene (25), was attributed to Homo, due, in part, to the small tooth crown dimensions (26), but was also regarded as morphologically compatible with P. robustus (27). More than a dozen specimens from Sterkfontein member 4 (~2.8 Ma to 2.2 Ma) and member 5 (2.2 Ma to 2.0 Ma) (4), have been attributed to Homo (SE 255, SE 1508, SE 1579, SE 1937, SE 2396, Sts 19, StW 19b, StW 42, StW 53, StW 75, StW 80 and 81, StW 84, StW 87, and StW 151) (1, 2, 16, 27), although some specimens, including StW 53, were also regarded as representative of A. africanus (28, 29). In addition to the recently described calvaria (6), 10 isolated teeth from Drimolen (DNH 24, DNH 35, DNH 39, DNH 42, DNH 45, DNH 62, DNH 67, DNH 70, DNH 71, and DNH 80) were also suggested to represent early Homo (30, 31).

Collectively, this southern African sample of the early Homo hypodigm exhibits a high degree of dentognathic dimensional and morphological variation, and the attribution to Homo of some specimens remains a matter of contention (e.g., refs. 1, 2, 6, 19, 20, 22, 24, 27, and 32-40). Indeed, this assemblage has been referred to as Homo sp., Homo habilis, and H. erectus/H. ergaster, but many of the specimens differ in dental morphology from the eastern African remains allocated to the latter two species (27). This casts doubts on the nature and identity of the southern African Early Pleistocene material regarded as Homo. In addition, a new species Homo gautengensis (40) was erected to accommodate most of the Early Pleistocene southern African Homo-like specimens. StW 53 was defined as the holotype, along with a number of paratypes (SE 255, SE 1508, StW 19b/33, StW 75-79, StW 80, StW 84, StW 151, SK 15, SK 27, SK 45, SK 847, SKX 257/ 258, SKX 267/268, SKX 339, SKX 610, SKW 3114, and DNH 70). However, the type specimen of H. gautengensis, StW 53, has been generally regarded as representing H. habilis (27), and has also been shown to be morphologically indistinguishable from A.

africanus (28, 29). For this reason, the validity of this endemic taxon has not been generally accepted, and the attribution of the specimens remains controversial (41). Disagreements partly arose because previous studies were restricted to the analysis of a few incomplete mandibles and cranial remains chronologically spanning 700 ka to 500 ka (27). However, the majority of the southern African *Homo* record is represented by dental remains, and the analysis of the internal structure of these teeth has the potential to shed light on their taxonomic affiliation, augmenting debate around the origins and distribution of early *Homo* in the Cradle of Humankind.

To reassess the taxonomic attribution of the southern African early Homo dentognathic assemblage, we focus on the morphology of the enamel-dentine junction (EDJ). This interface is preserved in fully formed tooth crowns between enamel and dentine and captures the morphology of the membrana praeformativa, a basement membrane serving as the template for the majority of the taxonomically relevant aspects of crown morphology (42, 43). Unlike the outer enamel surface, which is often affected by occlusal wear, the EDJ is generally well preserved and recognized as a reliable taxonomic proxy to distinguish between hominid taxa, even at the subspecies level (44-48). Three-dimensional geometric morphometric (3D GM) analyses using landmarks and semilandmarks show that dentine horn height, crown height, and cervix shape can distinguish the postcanine teeth of Australopithecus, P. robustus, and Late Pleistocene to Holocene Homo (47, 49, 50). The recently developed diffeomorphic surface matching (DSM) method (51), which models the deformation between shapes, captures the geometric details of an anatomical structure and is now used as an advanced analytical tool in morphometrics (51-54). Indeed, DSM analyses can capture the taxonomically relevant aspects of the EDJ morphology, including both prominent features (such as the dentine horns and marginal crests) and more subtle features (like the protostylid and occlusal basin morphology), and might thus improve on traditional GM analyses that focus only on the shape of the cervix and marginal ridge (53, 55, 56).

We use the DSM approach to investigate the EDJ shape of an assemblage of permanent premolars and molars from Drimolen (DNH 39, DNH 62, DNH 67, and DNH 70), Kromdraai (KB 5223), Sterkfontein (SE 255, SE 1508, Sts 9, StW 19b, StW 53, StW 80, StW 81, StW 87, StW 151, and StW 669), and Swartkrans (SK 15, SK 18a, SK 27, SK 96, SK 847, SKX 257, SKX 268, and SKX 21204) attributed to, or suggested to represent, Homo (SI Appendix, Supplementary Note 1 and Table S1). We test whether their EDJ morphology more closely approximates the condition of Early-Middle Pleistocene *Homo* from eastern Africa (n = 22/65) and Asia (n = 43/65) or the australopith (Australopithecus and Paranthropus from southern Africa) pattern (for the list of the comparative material, see SI Appendix, Table S2). The reference Homo sample includes specimens attributed to H. erectus/H. ergaster or early Homo and that are widely accepted to unambiguously belong to our genus during the early period of its evolution.

Results

As illustrated in the assemblage from Swartkrans (Fig. 1), for all tooth positions, there are aspects of EDJ morphology of the southern African purported *Homo* specimens that are more typical of *Australopithecus* and *Paranthropus* (*SI Appendix*, Figs. S1–S8). To quantitatively assess EDJ shape variation and reassess taxonomic attribution of the purported *Homo* teeth, we conducted DSM GM analyses of the EDJ using specimens of *Australopithecus*, *Paranthropus*, and *Homo* whose taxonomic



Fig. 1. The EDJ of the postcanine teeth of the purported *Homo* specimens from Swartkrans compared with those of Early Pleistocene *Homo* (KNM-ER 1590 M^1 and M^2 , Sangiran 4 M^3 , and KNM-ER 992 lower postcanine teeth), *Australopithecus* (Taung M^1 and M_1 , StW 183 M^2 , StW 128 M^3 , StW 498 P_3 , StW 104 P_4 , StW 133 M_2 , and StW 312 M_3), and *Paranthropus* (TM 1517 upper molars and P_3 , P_4 , M_1 , and M_3 , and SK 1 M_2). Specimens belonging to the same individual are enclosed by a plain line, and specimens that likely belong to the same individual are enclosed by a dotted line.

identity at genus level is well established and undisputed as reference samples. In all between-group principal component analyses (bgPCA) based on the deformation fields computed by DSM, the three hominin genera are well discriminated, despite a slight overlap between Australopithecus and Paranthropus for the M^3 and P_4 (Figs. 2 and 3). The cross-validated bgPCA (cv-bgPCA) confirm the general distinction of the three groups. In this case, while the australopiths overlap for the premolars and third molars, Homo is largely well distinguished (SI Appendix, Figs. S9 and S10). Additionally, we conducted canonical variate analyses (CVA) on subsets of the principal components, and results are consistent with those of the bgPCA and cv-bgPCA (SI Appendix, Figs. S11 and S12). As shown by the statistical analyses, all cv-bgPCA and CVA of PC scores exhibit high degrees of classification accuracy for the comparative specimens (SI Appendix, Tables S3 and S4), and group differences are not purely allometric, as they remain large and significant after controlling for allometry (SI Appendix, Table S5). With respect to the australopiths, *Homo* generally shows a taller EDJ crown with a smaller EDJ central basin in the premolars, and a proportionally larger basin with respect to the crown base (except in the third molars) and a rounder or

more regular cervical outline in the molars. *Australopithecus* is distinguished from *Paranthropus* by a more developed lingual than buccal aspect in upper molars, a less distally positioned protoconid in the P_3 , a shorter P_4 EDJ crown, and a more expanded buccal shelf in the lower molars.

The purported Homo specimens were projected into the bgPCA and CVA, and the typicality probabilities were computed (Table 1). With a few exceptions, the results of the bgPCA and CVA are consistent. Specimens from Drimolen (DNH 39, DNH 62, DNH 67, and DNH 70) and Kromdraai B (KB 5223) are statistically attributed to Paranthropus, sharing with the teeth of this genus a more mesiodistally compressed lingual aspect of the EDJ than the buccal one in the upper molars (Fig. 2) and a developed lower molar protostylid (Fig. 3). The specimen SK 15 also shows strong affinities with Paranthropus (of the three lower molars analyzed, the M₂ and left M₃ fall within *Paranthropus*, while the right M₃ is intermediate between Paranthropus and Australopithecus). The P_{3/4} SK 18a, that was suggested to belong to the SK 15 mandible, also displays an intermediate signal, and is closer to Australopithecus if it is a P₄ (Figs. 1 and 3). The specimens SE 255, StW 19b, StW 87, and StW 151 from Sterkfontein and SKX 257 and



Fig. 2. Bivariate plot of the bgPCA scores based on the DSM deformation fields for the M¹ (*A*), M² (*B*), and M³ (*C*). The totality of the variance refers to between-group variation. Symbols highlighted in bold represent the holotype specimens of *Australopithecus* (Taung) and *Paranthropus* (TM 1517). Filled triangles indicate African *Homo*, while open triangles represent Asian *Homo*.

SKX 268 from Swartkrans are distinctly classified within *Australopithecus* (even if the M^2 of StW 151 is closer to the *Paranthropus* morphology). The M^3 s of StW 53 exhibit a mosaic of *Paranthropus* and *Australopithecus* morphology, while the M₃ shape is compatible with the *Australopithecus* variation.

Conversely, the specimens SK 27 and SKX 21204 from Swartkrans, as well as Sts 9 from Sterkfontein, fall well within *Homo*, showing a taller EDJ crown than in the australopiths (Figs. 2 and 3). The M^3 of SK 847 has a mesiodistally compressed shape that is similar to that of *Homo*, but also a trapezoidal outline reminiscent of the *Australopithecus* condition. The classification of the specimens StW 80 and 81 from Sterkfontein varies depending on tooth position. The P₃ and M₃ EDJ are comparable to *Homo* but outside the variation range displayed by the comparative sample used in this study, while the P₄ exhibits an australopith morphology, and the M₂ is intermediate between *Homo* and *Australopithecus*. The specimens SK 96 from Swartkrans, as well as SE 1508 and StW 669 from Sterkfontein, are less definitive, as their EDJ shape is intermediate between all three genera (Figs. 2 and 3 and *SI Appendix*, Figs. S11 and S12).

We also scored nonmetric morphological features of the EDJ in the purported early *Homo* and comparative samples, showing that there is an overlap in the presence/absence and degree of expression of most nonmetric dental features recorded at the EDJ of the three hominin genera. None of these traits in isolation can be considered diagnostic of *Homo* (*SI Appendix, Supplementary Note 2* and Table S6).

Finally, we investigated whether tooth size (assessed here using the EDJ surface area as a proxy) is a diagnostic feature to distinguish early *Homo* from the australopiths. With the exceptions of SK 15, StW 19b, StW 81, and StW 53 that have relatively large EDJ areas (in the upper range of *Australopithecus* and *Paranthropus*), the size of most of the purported early *Homo* specimens is in the lower range of the australopiths and within the range of Early to Middle Pleistocene *Homo* (*SI Appendix*, Fig. S13). In addition, even if Early to Middle Pleistocene *Homo* tends to have smaller tooth dimensions than the australopiths, there is overlap with *Australopithecus* for most tooth positions. A summary of the results for the metric, nonmetric, and size analyses is presented in *SI Appendix*, Tables S6–S8 along with an overall assessment of the taxonomic affiliation of each of the purported *Homo* specimens.

Discussion

The definition of the genus *Homo* on paleontological grounds remains a conundrum (57, 58). When the species *H. habilis*

dental features distinguishing Homo from Australopithecus, including a bicuspid P₃, smaller postcanine teeth (in particular the M^3 , generally smaller than the M^2 or M^1), anterior teeth proportionally less reduced than the postcanine elements, and less buccolingually expanded premolars and molars (essentially due to the reduction or absence of the protostylid in Homo). However, these features are also variably found in australopith specimens, sometimes in combination (for example, the Australopithecus specimen Sts 52 has a smaller M³ relative to the M² and a bicuspid P₃), whereas some specimens attributed to Homo (e.g., the H. habilis specimen OH 16) show a larger M3 than M2 and a mesiodistally elongated P₃ (36, 59). Wood and Collard (57) suggested that fossil specimens/species should be included in Homo only if their teeth are more similar in morphology and relative proportions to the modern human condition than they are to the australopiths. However, modern human tooth morphology differs from most of the earlier species of *Homo*: The P_3 and P_4 have a small to absent metaconid and have a high crown, the M² and M³ variably display only three cusps (lacking a hypocone), and the M₂ and M₃ often have only four cusps (SI Appendix, Supplementary Note 2). This is largely due to the trend for reduction of the size and prognathism of the face, the jaw, and tooth dimensions that occurred through the Late Pleistocene (60). Conversely, Early to Middle Pleistocene Homo species have more-complex lower premolars (with a large metaconid and an extended talonid), four-cusped upper molars, and five-cusped lower molars, making them more suitable for comparison with the Late Pliocene to Early Pleistocene record and the identification of early Homo.

was erected, Leakey et al. (59) proposed several morphological

In the bgPCA, cv-bgPCA, and CVA of PC scores conducted here, fossil Homo, Australopithecus, and Paranthropus are generally well discriminated, further demonstrating the relevance of the EDJ for taxonomic diagnosis (44, 45, 48, 53). The Early to Middle Pleistocene reference sample includes both African and Asian specimens that group closely and are mixed together in the plots despite their wide chronogeographic distribution. Due to the inherent nature of the fossil record, hominin taxa (including early Homo species) are represented by a limited number of specimens, which complicates assessment of intrataxonomic and intertaxonomic variation, and, ultimately, taxonomic attribution (27, 57). However, in the absence of molecular data for early hominins, studying morphology remains the only way to diagnose extinct taxa. Among the 23 southern African specimens previously suggested to belong to Homo examined in this study (Table 2), only three are unequivocally attributed to Homo (SK 27, SKX 21204,



Fig. 3. Bivariate plot of the between-group principal component analysis (bgPCA) scores based on the DSM deformation fields for the P₃ (*A*), P₄ (*B*), M₁ (*C*), M₂ (*D*), and M₃ (*E*). The totality of the variance refers to between-group variation. Symbols highlighted in bold represent the holotype specimens of *Australopithecus* (Taung) and *Paranthropus* (TM 1517). Filled triangles indicate African *Homo*, while open triangles represent Asian *Homo*.

and Sts 9). SK 847 morphology is closer to *Homo*, even if the overall EDJ configuration retains *Australopithecus* features (Fig. 2 and *SI Appendix*, Fig. S3) and can also be confidently attributed to the former genus. This is compatible with the analysis of the morphology of the face and partial cranium of this specimen

showing that it shares synapomorphies of the *Homo* clade but probably represents a more primitive species than *H. erectus* s.l (18, 39). The specimens StW 80/81, representing a single individual, preserve crushed and fragmentary portions of the mandibular bone with little morphological information (1). The EDJ of

the postcanine teeth of StW 80/81 displays affinities with Homo, as well as some features reminiscent of the australopith condition (mesiodistally elongated P4 and M3, and developed protostylid on the M₂). This combination of Australopithecus-like and Homolike traits detected in some specimens investigated here can be interpreted in different ways: 1) These individuals could represent some of the earliest members of the genus Homo, retaining some primitive features of the dentition; 2) they could belong to an Australopithecus group evolving directly or convergently toward the Homo condition; or 3) they could be an australopith-Homo hybrid of the kind hypothesized by Thackeray (61). With respect to the first and second scenarios, StW 80/81 comes from member 5 west of Sterkfontein, likely dating to 1.7 Ma to 1.4 Ma (1). If this individual represents an early member of our genus with a morphology reminiscent of Australopithecus, it could imply that a stem group of early Homo remained genetically isolated for some time from H. erectus/H. ergaster. If it represents Australopithecus, it is more recent than A. sediba, and future studies of the EDJ of the postcanine dentition could resolve whether StW 80/81 morphology is compatible with A. sediba. The third hypothesis cannot be discarded either, and, while the influence of interbreeding on hominin tooth morphology is poorly understood, molecular analyses such as paleoproteomics should be soon able to test this and investigate phylogenetic relationships of Pliocene and Early Pleistocene hominins (62).

The case of the specimens SE 1508 and StW 669, from Sterkfontein, is more ambiguous, as they have a central position in the bgPCA and/or CVA plots (Fig. 2 and SI Appendix, Fig. S11), indicating the need for more caution in the interpretation of their group affiliation. The EDJ of the M² SE 1508 is similar to that of SK 27, albeit with a lower and broader crown base with respect to the occlusal basin. In the DSM analyses, these differences drive the specimen away from the Homo morphology represented in our sample and make it closer to Paranthropus. However, together with the marked similarities with SK 27, the position of the dentine horns and relatively simple crown morphology (i.e., absence of accessory traits) suggest that SE 1508 could represent early Homo. The M1 StW 669 shows external crown morphological features similar to those of *H. habilis* (63), and preliminary analyses of the internal tooth structure also show affinities with Homo (64). However, our comparative analyses reveal that the EDJ retains some Australopithecus-like features, such as a markedly waisted occlusal outline, a buccal shelf, and a mesial lateral aspect that is strongly oriented inwardly (as in StW 283, for example). While an attribution to early Homo cannot be rejected, the specimen may represent Australopithecus with derived features resembling the Homo condition.

All the other purported early *Homo* specimens analyzed here show a dominant australopith signature of the EDJ morphology, and statistical analyses suggest that their inclusion in the *Homo* hypodigm is poorly supported based on EDJ morphology. As many of these are isolated teeth, additional analyses will be necessary to more clearly establish their taxonomic status (e.g., proteomic and ancient DNA evidence, and/or isotopic data [see below]). This also implies that the species *H. gautengensis*, created to accommodate the large morphodimensional variation of the southern African "early *Homo*" material (40), is invalid, as it includes specimens belonging to *Homo* (e.g., SKX 21204 and SK 27), *Australopithecus* (e.g., SE 255 and StW 53), and *Paranthropus* (e.g., KB 5223 and DNH 70).

SK 96 has a relatively central position in the bgPCA (Fig. 3), while the CVA place it close to *Paranthropus* (*SI Appendix*, Fig.

Table 1. Typicality probabilities of the investigated specimens computed for the bgPCA and CVA of shape analyses

	НОМ		AUS		PAR	
	bgPCA	CVA	bgPCA	CVA	bgPCA	CVA
M ¹ DNH 39 DNH 62 DNH 70					0.19 0.34 0.37	0.25 0.08 0.11
SE 255 SK 27 L	0.16 0.18	0.42 0.55	0.82	0.18	0.57	0.11
SK 27 R SKX 268 StW 151 L StW 151 R StW 669			0.24 0.39 0.28 0.19	0.07 0.68 0.58 0.01		
M ² SE 1508	0.90	0.75			0.06	0.39
SK 27 StW 151 M ³	0.80	0.75			0.29	0.52
SK 847 StW 19 StW 53 L	0.22		0.50 0.46	0.02 0.30	0.24	0.09 0.26
StW 53 R P ₃					0.50	0.76
SK 18a SK 96 SKX 21204 StW 80	0.36 0.05	0.89 0.74	0.21		0.03	0.16 0.07
SK 18a	0.21	1 0.32	0.26	0.32		
StW 80 StW 87 StW 151 L StW 151 R	0.51		0.25 0.11 0.79 0.38	0.58 0.06 0.31 0.73		
M ₁ DNH 67 KB 5223					0.99 0.14	0.99 0.12
SKX 257 Sts 9	0.84	0.47	0.52	0.63		
StW 151 L StW 151 R			0.71 0.60	0.77 0.76		
M ₂ SK 15 StW 80 M ₂			0.19	0.62	0.81	0.53
SK 15 L SK 15 R StW 53h			0.31 0.37	0.42 0.01	0.61	0.10

For each specimen, typicality probabilities are shown only for the group with which that specimen was affiliated. Values in bold are below the 0.05 threshold and are regarded as outliers for the groups included here. AUS, *Australopithecus*; HOM, Early and Middle Pleistocene *Homo*; PAR, *Paranthropus*.

S12), as also indicated by the typicality probabilities (Table 1). The bgPCA probabilities are below the statistical threshold of classification within any of the comparative groups (P < 0.05), suggesting that SK 96 could either be an outlier for any of the comparative genera or represent another genus that is not represented in the reference samples. It was suggested by Davies et al. (50) that SK 96 EDJ shares similarities with *Paranthropus* and also approximates that of *Homo naledi*. The taxonomic

Specimen	Tooth	EDI area	Nonmetric traits	cv-bgPCA	cv-CVA	Bone morphology	Revised assessment
DNH 39	RM ¹		UND	PAR	PAR	n/a	PAR
DNH 62	I M ¹		UND	PAR	PAR	n/a	PAR
DNH 67	RM₄	AUPITH	UND	PAR	PAR	n/a	PAR
DNH 70	I M ¹	UND	UND	PAR	PAR	n/a	PAR
KB 5223	LM	AUPITH	AUPITH	PAR	PAR	n/a	PAR
SE 255	RM ¹	UND	UND	AUS	AUS	n/a	AUS
SE 1508	RM ²	НОМ	UND	INT	PAR	n/a	PAR?
StW 19b	RM ³	AUPITH	UND	AUS	AUPITH	n/a	AUS
StW 53	LM ³	AUPITH	UND	AUPITH	PAR	HOM?	AUPITH
StW 53	RM ³	AUPITH	UND	AUPITH	PAR	HOM?	AUPITH
StW 53	LM3	AUPITH	UND	AUS	AUS	HOM?	AUS
StW 80	RP ₃	UND	UND	НОМ	НОМ	UND	НОМ
StW 80	RP₄	HOM?	UND	AUS?	AUS	UND	AUS/HOM
StW 80	RM_2	UND	UND	AUS?	AUS	UND	AUS?
StW 81	RM3	AUPITH	UND	INT	НОМ	UND	НОМ
StW 87	RP₄	HOM?	UND	AUS?	AUS	n/a	AUS
StW 151	LM ¹	UND	UND	AUS	AUS	UND	AUS
StW 151	RM ¹	UND	UND	AUS	AUS	UND	AUS
StW 151	RM ²	НОМ	UND	PAR	PAR	UND	AUPITH
StW 151	LP₄	НОМ	UND	AUS	AUS	UND	AUS
StW 151	RP₄	НОМ	UND	AUS?	AUS	UND	AUS
StW 151	LM ₁	UND	AUPITH?	AUS	AUS	UND	AUS
StW 151	RM₁	UND	AUPITH?	AUS	AUS?	UND	AUS
StW 669	RM ¹	UND	UND	AUS?	AUS	n/a	AUS?
Sts 9	RM_1	AUPITH	UND	НОМ	НОМ	n/a	НОМ
SK 15	RM ₂	AUPITH	UND	PAR	PAR	HOM?	PAR
SK 15	LM ₃	AUPITH	UND	PAR	PAR	HOM?	PAR
SK 15	RM ₃	AUPITH	UND	AUS?	AUS?	HOM?	AUPITH
SK 18a	LP_3 or LP_4	HOM?	UND	AUS	AUPITH	HOM?	AUPITH
SK 27	LM ¹	UND	UND	ном	ном	НОМ	ном
SK 27	RM ¹	UND	UND	HOM?	ном	НОМ	ном
SK 27	RM ²	ном	UND	ном	ном	НОМ	ном
SK 96	LP ₃	UND	UND	AUPITH	PAR?	n/a	AUPITH
SK 847	LM ³	AUPITH	UND	ном	AUS?	НОМ	ном
SKX 257	RM_1	UND	UND	AUS	AUS	n/a	AUPITH
SKX 268	RM ¹	UND	AUPITH	AUS	AUS	n/a	AUPITH
SKX 21204	RP₃	НОМ	UND	ном	ном	ном	ном
SKX 21204	RP ₄	НОМ	UND	НОМ	НОМ	НОМ	НОМ

Table 2. Summary of taxonomic evidence for specimens previously attributed to Homo

Bold and nonbold rows separate the different individuals represented here. AUS, Australopithecus; AUPITH, australopith; HOM, Early and Middle Pleistocene Homo; INT, intermediate; PAR, Paranthropus; UND, undiagnostic.

attribution of this specimen thus remains problematic. If SK 96 belongs to Paranthropus, it differs from the morphology of P. robustus, and it raises questions regarding possible affinities with *H. naledi*. There is a chronological gap of more than 1 My between the Swartkrans member 2 specimen and the Rising Star Cave hominins, and phylogenetic relationships of H. naledi with other hominin taxa are still uncertain (50, 65), and could be a focus of future studies. Our results also indicate that SK 15, holotype of the nomen oblitum T. capensis (15), as well as SK 18a that is supposedly associated with this mandible, actually represent an australopith taxon and not Homo. Although generally regarded as a representative of H. erectus s.l (9), the specimen is extremely robust, with proportions of the mandibular corpus that markedly differ from those of H. erectus/H. ergaster and other Pleistocene Homo species (66). The EDJ of SK 15 molars shows strong affinities with Paranthropus but differs in size and in relevant morphological aspects from the species P. robustus (e.g., absence of protostylid and accessory features) (Fig. 1). Interestingly, SK 15 molar root morphology has been linked with H. naledi and is, to some extent, also comparable with the

Paranthropus condition (67). If SK 15 belongs to *P. robustus*, then it either suggests high levels of variation in mandible size/shape with implications for sexual dimorphism within the genus or a higher degree of taxonomic diversity than currently recognized in the genus. The currently available evidence from dental structural organization suggests that SK 15 possibly belongs to *Paranthropus*, but its taxonomic status should be investigated further.

Based on morphology and dimensions of the craniodental remains, the specimens StW 53 and StW 151 were both suggested to represent early *Homo* or *Australopithecus* individuals more derived toward *Homo* than the rest of the australopith hypodigm (2, 28, 29, 68). StW 53 preserves partial aspects of the face and cranium (69). The most recent reconstruction shows a small endocranial volume and an intermediate morphology between *Australopithecus* and *H. habilis* (37), even if it has been criticized for the anatomically unrealistic shape, indicating that the reconstruction of the cranium would be closer to *Australopithecus* (29). The juvenile specimen StW 151 only preserves small bone fragments of the jaw and cranium that are not very informative for taxonomy (68). Our results show



Fig. 4. Chronostratigraphic distribution of the investigated purported *Homo* remains. In light of the results of the present study, only four specimens are likely to represent *Homo* (green ticks), four are possibly *Homo* (light green question marks), and the others more likely belong to *Australopithecus* or *Paranthropus* (red crosses).

that EDJ morphology of both specimens is on the margins of *Australopithecus* variation or even approximates the *Paranthropus* condition. Interestingly, the analysis of bony labyrinth shape shows that the two specimens represent the two extremes of the *Australopithecus* variation, where StW 53 appears as an outlier of the taxon (70).

All the Drimolen specimens investigated here that were suggested to represent early Homo (DNH 39, DNH 62, DNH 67, and DNH 70) show EDJ morphology that is more similar to P. robustus. Interestingly, the M₁ KB 5223, whose crown shows small dimensions comparable to those of early Homo (26) and smaller than in *P. robustus*, likely belongs to the latter taxon. Indeed, the EDJ of this specimen displays a marked protostylid that is similar to that of P. robustus. It is noteworthy that the analysis of enamel microstructure of KB 5223 revealed an overall pattern compatible with Paranthropus (71). The above specimens from Drimolen are relatively small and are dated to around 2 Ma (6), and KB 5223 from Kromdraai comes from early Early Pleistocene sediments (25), again suggesting that a smaller morph of P. robustus than that from Swartkrans coexisted in southern Africa for some time. However, the chronology of the deposits from these sites is debated, and the small Paranthropus dental specimens from Drimolen and Kromdraai that were previously identified as Homo could represent the early condition of the P. robustus morphocline (72, 73).

Most of the Early Pleistocene southern African teeth that our analyses identify as misattributed to Homo are smaller than those typical of Australopithecus and Paranthropus, while those compatible with *Homo* in shape are relatively large (SI Appendix, Fig. S13). It is possible that, besides the smaller dimensions, the external crown shape initially used to classify these teeth is also less diagnostic. In this respect, allometric changes could be considered as a potential factor. In our PCA, bgPCA, and CVA, allometric changes could be considered as a potential factor. However, even after controlling for allometry, group differences remain large (R² ranging from 9 to 29%) and significant (SI Appendix, Table S5). This suggests that differences between EDJ morphology of the specimens mostly represent shape variation and are only modestly influenced by size. This study also demonstrates that, at many postcanine tooth positions, crown dimensions overlap in Australopithecus, Homo, and Paranthropus (SI Appendix, Fig. S13). Using the natural logarithm of the EDJ as a proxy for crown size, Early to Middle Pleistocene Homo generally has significantly smaller postcanine teeth than *Paranthropus* (except for the M¹) and slightly smaller

teeth than *Australopithecus*, on average (but only partially, as indicated by the small determination coefficient and *SI Appendix*, Fig. S13 and Table S7). In fact, some of the investigated specimens show small dimensions, close to those of *Homo*, but the EDJ shape unambiguously displays an australopith morphology (e.g., DNH 70 and StW 151), and vice versa (SK 27 and SK 847 have large molars but a *Homo*-like EDJ shape). This result thus indicates that tooth size alone should not be used for taxonomic diagnosis in this lineage.

Following our revision, among the 23 specimens previously labeled as Homo, only between 4 and 7 specimens very likely belong to this taxon. It is possible that some of the southern African purported Homo specimens that, in our analysis, exhibit an australopith-like signal actually do belong to Homo. This would mean that, dentally speaking, the typical tooth morphology of specimens attributed as H. erectus/H. egaster around 2 Ma did not emerge with the genus or was not ubiquitous among all early Homo groups. However, the diversity of Australopithecus-like and Paranthropus-like signals that we have found in the southern African purported Homo hypodigm suggests that it is unlikely that all of these specimens belong to Homo. We propose that, in addition to the four that display a Homo signal, a number of other individuals that show australopith affinities (e.g., SE 1508, StW 80 and 81, and StW 669) should be investigated further and compared with the hominin specimens from Olduvai Gorge, Tanzania, currently under study for internal tooth structure (74). The presence of another hominin taxon not included in the reference sample and represented by one or more of these intermediate specimens cannot be discarded either. This revision of the purported early Homo dental material reduces its frequency and representation in the Early Pleistocene southern African fossil record (Fig. 4), and this has implications for the interpretation of the fossil hominin paleodiversity, paleoecology, and dietary behavior at macroregional scale (75).

As an example, if a specimen is regarded as *Homo* whereas it belongs to *Australopithecus* or *Paranthropus*, the interpretations of the biological signal that can be extracted from the mineralized tissues are erroneous. This leads to the false impression that *Homo* was more variable in growth, development, morphology, behavior, and ecology than it really was, and prevents understanding of how it biologically differed from other hominin genera. To illustrate this, we analyzed here the geochemical profile of three teeth previously regarded as *Homo*: SK 96, whose taxonomic identity remains uncertain, and KB 5223 and SKX 268, reattributed, in this study, to *Paranthropus* and *Australopithecus*,



Fig. 5. Ba/Ca (A) and Sr/CA (B) ratios and elemental mapping of SK 96, KB 5223, and SKX 268 compared with the distributions of *Australopithecus* and *Paranthropus*. Asterisks denote that data for taxa distribution were calculated using a combination of new data and a previously published datasets (80, 81), with the average values of KB 5223 (triangles) and SKX 268 (crosses) extracted from Balter et al. (76). SK 96 profile analysis was realized on the enamel for this study, as well as the geochemical maps of KB 5223 and SKX 268. All error bars are 2 sigma.

respectively (Fig. 5). We also conducted elemental mapping of Sr/Ca and Ba/Ca ratios on the latter two specimens. Our geochemical results confirm those of a previous analysis of KB 5223 and SKX 268 that aimed to infer early Homo dietary preferences (76). Both SK 96 and KB 5223 have low Sr/Ca ratios, close to the mean value of TM 1517, holotype of P. robustus, with a range of variation compatible with the values of penecontemporaneous browser taxa (76). The elemental maps of KB 5223 are highly affected by diagenesis, showing large areas of uniform density and few structured areas indicative of a biogenic signal (i.e., of biological origin, as opposed to diagenetic that is related to taphonomy and fossilization processes). In the M1 SKX 268, the Ba/Ca and Sr/Ca ratios are higher than in the other two specimens, more closely approximating the condition of the Australopithecus. The overall distribution of Ba/Ca in the enamel indicates a dominance of biogenic signal (some of the highest concentrations follow the striation pattern of the Retzius lines), while the Sr/Ca ratio is more influenced by diagenetic processes (with low and high values distributed homogeneously). Conversely, for both variables, the dentine shows pristine, biogenic banding typical of the growth

layering architecture of tooth formation, mimicking the profiles obtained by a previous study on *Australopithecus* (77). SKX 268 preserves part of the nursing sequence, which is comparable with the breastfeeding timing previously proposed for *Australopithecus* (77). Lactation behavior—with the infant relying more on solid food during periods of abundance, allowing the mother to replenish her energetic and calcium reserves to support lactation during periods of food scarcity—was proposed to represent a versatile adaptive trait to survive challenging ecological niches (77). Taxonomically misattributing SKX 268 would thus skew our understanding of the intrataxic/intertaxic variability of this feature.

Regardless of the robustness of the biogeochemical studies arising from the southern African hominin fossil record, accurate interpretations are contingent on correct taxonomic classification of hominin dental specimens. Our results highlight the need to continually interrogate taxonomic attributions of Early Pleistocene hominins in light of new evidence, such as shifts in first appearance dates [i.e., necessitated by the classification of the Ledi-Geraru mandible as Homo (78)], the unresolved taxonomic status of numerous specimens from Omo, Ethiopia (79), and Koobi Fora, Kenya (36), and the recent discovery of rather primitive Homo dentitions from Rising Star Cave, South Africa (65), Liang Bua, Indonesia (80), and Callao Cave, Philippines (81). This will facilitate paleobiological reconstructions and our understanding of the timing, location, and nature of the selective pressures that resulted in a shift between australopith-like and Homo-like grades during hominin evolution.

Materials and Methods

Thirty-seven southern African postcanine teeth previously attributed to early *Homo*, as well as one isolated lower molar, Sts 9, generally attributed to *Australopithecus* but recognized here as *Homo* (*SI Appendix*, Table S1), were compared with *Australopithecus* (n = 123), *Paranthropus* (n = 97), and African and Asian Early to mid-Middle Pleistocene *Homo* (n = 66; *SI Appendix*, *Supplementary Note 1* and Table S2). We combined nonmetric trait and DSM GM approaches to investigate the EDJ shape of the purported early *Homo* sample, and we conducted geochemical analyses on three of the investigated specimens (for more detailed information, see *SI Appendix*, *Supplementary Material and Methods*).

Data Availability. All study data are included in the article and/or *SI Appendix*. The microCT scans of fossil specimens are available by research application to the relevant curatorial institution.

ACKNOWLEDGMENTS. For their contribution with scanning, granting access to, and/or sharing data of some of the fossil hominin specimens included in this study, we acknowledge J. W. Adams, C. Argot, F. Bernardini, J. Braga, L. Fiorenza, D. Grimaud-Hervé, J. Kibii, H. Lelièvre, L. Mancini, A. Mazurier, E. Mbua, S. Potze, F. Sémah, C. Tuniz, and H. Widianto, as well as the online platform European Synchrotron Radiation Facility heritage database for paleontology, evolutionary biology, and archaeology (paleo.esrf.eu). For scientific discussion and comments, we thank R. J. Clarke, F. Détroit, J. Dumoncel, F. E. Grine, S. Schlager, A. Urciuoli, and B. A. Wood. For technical support and access to tomographic scans and associated digital data, we thank the following institutions: University of the Witwatersrand (Johannesburg), Ditsong National Museum of Natural History (Pretoria), Max Planck Institute for Evolutionary Anthropology (Leipzig), Centre de Microtomographie Université de Poitiers, South African Nuclear Energy Corporation SOC Ltd. (Pelindaba), and International Centre for Theoretical Physics (Trieste). We thank the South African Heritage Resources Agency for authorizing the study of South African fossil material. We acknowledge support from the National Center for Scientific Research (CNRS)/IN2P3 Computing Center (Lyon, France) for providing computing and data processing resources needed for this work. We thank the editor-in-chief, the editor, A. Cardini, R. J. Clarke, and two anonymous reviewers for their constructive comments that helped to improve previous versions of this article. This research was funded by Erasmus+ Programme of the European Union Bakeng

se Afrika, French CNRS, AESOP+ program; National Research Foundation (South Africa); and Max Planck Society. This research benefited from the scientific framework of the University of Bordeaux's IdEx "Investments for the Future" program/ GPR "Human Past." The participation of M.M.S. was supported by funding from the European Research Council under the European Union's Horizon 2020 research and innovation program (Grant Agreement 819960).

Author affiliations: ^aUniv. Bordeaux, CNRS, MCC, PACEA, UMR 5199, F-33600 Pessac, France; ^bDepartment of Human Evolution, Max Planck Institute for Evolutionary Anthropology, 04103 Leipzig, Germany; ^cSchool of Anthropology and Conservation, University of Kent, Canterbury CT2 7NZ, United Kingdom; ^dGeoarchaeology and Archaeometry Research Group, Southern Cross GeoScience, Southern Cross University, Lismore, NSW 2480, Australia; ^ePalaeo-Research Institute, University of Johannesburg, Auckland Park 2006, South Africa; ^fDepartment of Archaeology, University

- K. Kuman, R. J. Clarke, Stratigraphy, artefact industries and hominid associations for Sterkfontein, member 5. J. Hum. Evol. 38, 827–847 (2000).
- J. Moggi-Cecchi, F. E. Grine, P. V. Tobias, Early hominid dental remains from Members 4 and 5 of the Sterkfontein Formation (1966-1996 excavations): Catalogue, individual associations, morphological descriptions and initial metrical analysis. J. Hum. Evol. 50, 239–328 (2006).
- A. I. R. Herries et al., "A multi-disciplinary perspective on the age of Australopithecus in Southern Africa" in *The Paleobiology of Australopithecus*, K. E. Reed, J. G. Fleagle, R. E. Leakey, Eds. (Springer, Dordrecht, 2013), pp. 21–40.
- D. E. Granger et al., New cosmogenic burial ages for Sterkfontein Member 2 Australopithecus and Member 5 Oldowan. Nature 522, 85–88 (2015).
- R. Pickering et al., U-Pb-dated flowstones restrict South African early hominin record to dry climate phases. Nature 565, 226–229 (2019).
- A. I. R. Herries et al., Contemporaneity of Australopithecus, Paranthropus, and early Homo erectus in South Africa. Science 368, eaaw7293 (2020).
- K. Kuman et al., A new absolute date from Swartkrans Cave for the oldest occurrences of Paranthropus robustus and Oldowan stone tools in South Africa. J. Hum. Evol. 156, 103000 (2021).
- 8. R. A. Dart, Australopithecus africanus: The man-ape of South Africa. Nature 115, 195-199 (1925).
- 9. B. A. Wood, Wiley-Blackwell Encyclopedia of Human Evolution (Blackwell, Chichester, 2011).
- R. A. Dart, The Makapansgat proto-human Australopithecus prometheus. Am. J. Phys. Anthropol. 6, 259-283 (1948).
- R. J. Clarke, K. Kuman, The skull of StW 573, a 3.67 Ma Australopithecus prometheus skeleton from Sterkfontein Caves, South Africa. J. Hum. Evol. 134, 102634 (2019).
- L. R. Berger et al., Australopithecus sediba: A new species of Homo-like australopith from South Africa. Science 328, 195–204 (2010).
- 13. R. Broom, Pleistocene anthropoid apes of South Africa. *Nature* **142**, 377–379 (1938).
- V. Balter et al., U-Pb dating of fossil enamel from the Swartkrans Pleistocene hominid site, South Africa. Earth Planet. Sci. Lett. 267, 236-246 (2008).
- 15. R. Broom, J. T. Robinson, A new type of fossil man. Nature 164, 322-323 (1949).
- J. T. Robinson, The Australopithecines and their bearing on the origin of Man and of stone toolmaking. S. Afr. J. Sci. 57, 3–13 (1961).
- R. J. Clarke, "Homo habilis: The inside story" in Proceedings of the II Meeting of African Prehistory, M. Sahnouni, S. Semaw, J. R. Garaizar, Eds. (Centro Nacional de Investigación Sobre la Evolución Humana, 2017), pp. 25–51.
- R. J. Clarke, F. C. Howell, C. K. Brain, More evidence of an advanced hominid at Swartkrans. *Nature* 225, 1219–1222 (1970).
- R. J. Clarke, A juvenile cranium and some adult teeth of early Homo from Swartkrans, Transvaal. S. Afr. J. Sci. 73, 46–49 (1977).
- F. E. Grine, New hominid fossils from the Swartkrans formation (1979-1986 excavations): Craniodental specimens. Am. J. Phys. Anthropol. 79, 409–449 (1989).
- T. R. Olson, Hominid phylogenetics and the existence of Homo in Member I of the Swartkrans Formation, South Africa. J. Hum. Evol. 7, 159-178 (1978).
- 22. J. H. Schwartz, I. Tattersall, Craniodental Morphology of Genus Homo (Africa and Asia) (The Human Fossil Record, Wiley-Liss, New York, 2003), vol. 2.
- M. C. Dean, B. A. Wood, Basicranial anatomy of Plio-Pleistocene hominids from East and South Africa. Am. J. Phys. Anthropol. 59, 157–174 (1982).
- F. E. Grine, Early Homo at Swartkrans, South Africa: A review of the evidence and an evaluation of recently proposed morphs. S. Afr. J. Sci. 101, 43–52 (2005).
- L. Bruxelles, R. Marie, R. Couzens, J. F. Thackeray, J. Braga, "A revised stratigraphy of Kromdraai" in Kromdraai: A Birthplace of Paranthropus in the Cradle of Humankind, J. Braga, Ed. (Sun, Stellenbosch, 2019), pp. 31–48.
- J. Braga, J. F. Thackeray, Early Homo at Kromdraai B: Probabilistic and morphological analysis of the lower dentition. C. R. Palevol 2, 269–279 (2003).
- F. E. Grine, H. F. Smith, C. P. Heesy, E. J. Smith, "Phenetic affinities of Plio-Pleistocene Homo fossils from South Africa: Molar cusp proportions" in *The First Humans - Origin and Early Evolution of the Genus Homo*, F. E. Grine, J. G. Fleagle, R. E. Leakey, Eds. (Springer, New York, 2009), pp. 49-62.
- W. W. Ferguson, Reappraisal of the taxonomic status of the cranium Stw 53 from the Plio/ Pleistocene of Sterkfontein, in South Africa. *Primates* 30, 103–109 (1989).
- R. J. Clarke, Latest information on Sterkfontein's Australopithecus skeleton and a new look at Australopithecus. S. Afr. J. Sci. 104, 443–449 (2008).
- A. W. Keyser, C. G. Menter, J. Moggi-Cecchi, T. R. Pickering, L. R. Berger, Drimolen: A new hominid-bearing site in Gauteng, South Africa. S. Afr. J. Sci. 96, 193–197 (2000).
- J. Moggi-Cecchi, C. Menter, S. Boccone, A. Keyser, Early hominin dental remains from the Plio-Pleistocene site of Drimolen, South Africa. J. Hum. Evol. 58, 374–405 (2010).
- F. C. Howell, "Hominidae" in *Evolution of African Mammals*, V. J. Maglio, H. B. S. Cooke, Eds. (Harvard University Press, Cambridge, 1978), pp. 154–248.
- F. E. Grine, "Implications of morphological diversity in early Homo crania from eastern and southern Africa" in *Humanity from the African Naissance to the Coming Millennia*, P. V. Tobias, M. Raath, J. Moggi-Cecchi, G. Doyle, Eds. (Firenza University Press, Firenza, 2001), pp. 107–115.

of Cambridge, Cambridge CB2 1TN, United Kingdom; ⁸School of Geography, Archaeology and Environmental Studies, University of the Witwatersrand, Johannesburg 2000, South Africa; ^hInstitut Català de Paleontologia Miquel Crusafont, Universitat Autònoma de Barcelona, 08193 Bellaterra, Spain; ¹UMR 5608 CNRS, TRACES, Maison de la Recherche, Université Toulouse Jean Jaurès, 31058 Toulouse Cedex 9, France; ¹INRAP, French Institute for Preventive Archaeological Researches, 30900 Nîmes, France; ¹South African Nuclear Energy Corporation SOC Ltd., Pretoria 0001, South Africa; ¹Department of Anthropology & Development Studies, University of Johannesburg, Auckland Park 2006, South Africa; ¹Evolutionary Studies Institute, University of the Witwatersrand, Johannesburg 2000, South Africa; ¹Ditsong National Museum of Natural History, Pretoria 0001, South Africa; ⁰Department of Paleoanthropology, Senckenberg Research Institute and Natural History Museum Frankfurt, Frankfurt 60325, Germany; ⁹Department of Paleobiology and Environment, Institute of Ecology, Evolution, and Diversity, Goethe University Frankfurt, 60323 Frankfurt, Germany; ⁹UMR 7194 CNRS, Muséum National d'Histoire Naturelle, 75005 Paris, France; ¹Département Géosciences, Université de Politiers, 86000 Poitiers, France; ^{*}Key Laboratory of Vertebrate Evolution and Human Origins, Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, 100044 Beijing, China; ^{*}Center for Excellence in Life and Paleoenvironment, Chinese Academy of Sciences, 100044 Beijing, China; and ^{*}Centre for the Exploration of the Deep Human Journey, University of the Witwatersrand, Johannesburg 2000, South Africa

- 34. G. P. Rightmire, The Evolution of Homo erectus (Cambridge University Press, Cambridge, 1990).
- P. V. Tobias, The Skulls, Endocasts and Teeth of Homo habilis (Olduvai Gorge, Cambridge University Press, Cambridge, 1991), vol. 4.
- 36. B. A. Wood, Hominid Cranial Remains (Koobi Fora Research Project, Clarendon, Oxford, 1991), vol. 4.
- W. H. Kimbel, Y. Rak, "The importance of species taxa in paleoanthropology and an argument for the phylogenetic concept of the species category" in *Species, Species Concepts, and Primate Evolution*, W. H. Kimbel, L. B. Martin, Eds. (Plenum, New York, 1993), pp. 461–485.
- D. Curnoe, P. V. Tobias, Description, new reconstruction, comparative anatomy, and classification of the Sterkfontein Stw 53 cranium, with discussions about the taxonomy of other southern African early *Homo* remains. J. Hum. Evol. 50, 36–77 (2006).
- H. F. Smith, F. E. Grine, Cladistic analysis of early *Homo* crania from Swartkrans and Sterkfontein, South Africa. J. Hum. Evol. 54, 684–704 (2008).
- D. Curnoe, A review of early *Homo* in southern Africa focusing on cranial, mandibular and dental remains, with the description of a new species (*Homo gautengensis* sp. nov.). *Homo* 61, 151–177 (2010).
- L. Berger, Australopithecus sediba and the earliest origins of the genus Homo. J. Anthropol. Sci. 90, 117–131 (2012).
- 42. P. M. Butler, The ontogeny of molar pattern. Biol. Rev. Camb. Philos. Soc. 31, 30-70 (1956).
- R. S. Corruccini, The dentinoenamel junction in primates. *Int. J. Primatol.* 8, 99–114 (1987).
 M. M. Skinner *et al.*, Dental trait expression at the enamel-dentine junction of lower molars in
- extant and fossil hominoids. J. Hum. Evol. 54, 173-186 (2008).
- M. M. Skinner, P. Gunz, B. A. Wood, C. Boesch, J.-J. Hublin, Discrimination of extant *Pan* species and subspecies using the enamel-dentine junction morphology of lower molars. *Am. J. Phys. Anthropol.* **140**, 234–243 (2009).
- R. Macchiarelli, P. Bayle, L. Bondioli, A. Mazurier, C. Zanolli, "From outer to inner structural morphology in dental anthropology. The integration of the third dimension in the visualization and quantitative analysis of fossil remains" in *Anthropological Perspectives on Tooth Morphology: Genetics, Evolution, Variation, R. G. Scott, J. D. Irish, Eds.* (Cambridge University Press, Cambridge, 2013), pp. 250–277.
- L. Pan et al., Intra-individual metameric variation expressed at the enamel-dentine junction of lower post-canine dentition of South African fossil hominins and modern humans. Am. J. Phys. Anthropol. 163, 806–815 (2017).
- C. Zanolli *et al.*, Evidence for increased hominid diversity in the Early to Middle Pleistocene of Indonesia. *Nat. Ecol. Evol.* **3**, 755–764 (2019).
- L. Pan et al., Further morphological evidence on South African earliest Homo lower postcanine dentition: Enamel thickness and enamel dentine junction. J. Hum. Evol. 96, 82–96 (2016).
- T. W. Davies et al., Distinct mandibular premolar crown morphology in Homo naledi and its implications for the evolution of Homo species in southern Africa. Sci. Rep. 10, 13196 (2020).
- S. Durrleman, X. Pennec, A. Trouvé, N. Ayache, J. Braga, Comparison of the endocranial ontogenies between chimpanzees and bonobos via temporal regression and spatiotemporal registration. J. Hum. Evol. 62, 74–88 (2012).
- A. Beaudet et al., Morphoarchitectural variation in South African fossil cercopithecoid endocasts. J. Hum. Evol. 101, 65–78 (2016).
- C. Zanolli *et al.*, Inner tooth morphology of *Homo erectus* from Zhoukoudian. New evidence from an old collection housed at Uppsala University, Sweden. *J. Hum. Evol.* **116**, 1–13 (2018).
- A. Urciuoli et al., Reassessment of the phylogenetic relationships of the late Miocene apes Hispanopithecus and Rudapithecus based on vestibular morphology. Proc. Natl. Acad. Sci. U.S.A. 118, e2015215118 (2021).
- J. Braga et al., Efficacy of diffeomorphic surface matching and 3D geometric morphometrics for taxonomic discrimination of Early Pleistocene hominin mandibular molars. J. Hum. Evol. 130, 21–35 (2019).
- L. Pan, J. Dumoncel, A. Mazurier, C. Zanolli, Hominin diversity in East Asia during the Middle Pleistocene: A premolar endostructural perspective. J. Hum. Evol. 148, 102888 (2020).
- 57. B. Wood, M. Collard, The human genus. Science 284, 65-71 (1999).
- B. Villmoare, Early *Homo* and the role of the genus in paleoanthropology. *Am. J. Phys. Anthropol.* 165, 72–89 (2018).
- L. S. B. Leakey, P. V. Tobias, J. R. Napier, A new species of the genus *Homo* from Olduvai Gorge. *Nature* 202, 7–9 (1964).
- C. L. Brace, "Cultural factors in the evolution of the human dentition" in *Culture and the Evolution of Man*, M. F. A. Montagu, Ed. (Oxford University Press, New York, 1962), pp. 343–354.
- J. F. Thackeray, Alpha and sigma taxonomy of Pan (chimpanzees) and Plio-Pleistocene hominin species. S. Afr. J. Sci. 114, 1-2 (2018).
- 62. F. Welker et al., The dental proteome of Homo antecessor. Nature 580, 235–238 (2020).
- D. Stratford, J. L. Heaton, T. R. Pickering, M. V. Caruana, K. Shadrach, First hominin fossils from Milner Hall, Sterkfontein, South Africa. J. Hum. Evol. 91, 167–173 (2016).
- B. Mataboge, A. Beaudet, J. L. Heaton, T. R. Pickering, D. Stratford, Endostructural assessment of a hominin maxillary molar (StW 669) from Milner Hall, Sterkfontein, South Africa. S. Afr. J. Sci. 115, 6404 (2019).

- L. R. Berger et al., Homo naledi, a new species of the genus Homo from the Dinaledi Chamber, South Africa. eLife 4, e09560 (2015).
- 66. D. Curnoe, Affinities of the Swartkrans early *Homo* mandibles. *Homo* **59**, 123–147 (2008).
- K. Kupczik, L. K. Delezene, M. M. Skinner, Mandibular molar root and pulp cavity morphology in Homo naledi and other Plio-Pleistocene hominins. J. Hum. Evol. 130, 83–95 (2019).
- J. Moggi-Cecchi, P. V. Tobias, A. D. Beynon, The mixed dentition and associated skull fragments of a juvenile fossil hominid from Sterkfontein, South Africa. Am. J. Phys. Anthropol. 106, 425–465 (1998).
- P. V. Tobias, Australopithecus, Homo habilis, tool-using and tool-making. S. Afr. Archaeol. Bull. 20, 167–192 (1965).
- A. Beaudet *et al.*, The bony labyrinth of StW 573 ("Little Foot"): Implications for early hominin evolution and paleobiology. *J. Hum. Evol.* **127**, 67-80 (2019).
- R. S. Lacruz, Enamel microstructure of the hominid KB 5223 from Kromdraai, South Africa. Am. J. Phys. Anthropol. 132, 175–182 (2007).
- J. M. Martin et al., Drimolen cranium DNH 155 documents microevolution in an early hominin species. Nat. Ecol. Evol. 5, 38-45 (2021).
- Y. Rak, W. H. Kimbel, J. Moggi-Cecchi, C. A. Lockwood, C. Menter, The DNH 7 skull of Australopithecus robustus from Drimolen (Main Quarry), South Africa. J. Hum. Evol. 151, 102913 (2021).

- 74. T. W. Davies *et al.*, Accessory cusp expression at the enamel-dentine junction of hominin mandibular molars. *PeerJ* **9**, e11415 (2021).
- M. Sponheimer, J. A. Lee-Thorp, Enamel diagenesis at South African Australopith sites: Implications for paleoecological reconstruction with trace elements. *Geochim. Cosmochim. Acta* 70, 1644–1654 (2006).
- V. Balter, J. Braga, P. Télouk, J. F. Thackeray, Evidence for dietary change but not landscape use in South African early hominins. *Nature* 489, 558–560 (2012).
- R. Joannes-Boyau *et al.*, Elemental signatures of Australopithecus africanus teeth reveal seasonal dietary stress. Nature 572, 112-115 (2019).
- B. Villmoare et al., Paleoanthropology. Early Homo at 2.8 Ma from Ledi-Geraru, Afar, Ethiopia. Science 347, 1352-1355 (2015).
- G. Suwa, T. D. White, F. C. Howell, Mandibular postcanine dentition from the Shungura Formation, Ethiopia: Crown morphology, taxonomic allocations, and Plio-Pleistocene hominid evolution. *Am. J. Phys. Anthropol.* **101**, 247–282 (1996).
- P. Brown *et al.*, A new small-bodied hominin from the Late Pleistocene of Flores, Indonesia. *Nature* 431, 1055–1061 (2004).
- F. Détroit *et al.*, A new species of *Homo* from the Late Pleistocene of the Philippines. *Nature* 568, 181–186 (2019).